This Page Is Inserted by IFW Operations and is not a part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images may include (but are not limited to):

- BLACK BORDERS
- TEXT CUT OFF AT TOP, BOTTOM OR SIDES
- FADED TEXT
- ILLEGIBLE TEXT
- SKEWED/SLANTED IMAGES
- COLORED PHOTOS
- BLACK OR VERY BLACK AND WHITE DARK PHOTOS
- GRAY SCALE DOCUMENTS

IMAGES ARE BEST AVAILABLE COPY.

As rescanning documents will not correct images, please do not report the images to the Image Problem Mailbox.

ISOTOPOME	R ABSORPTION SPECTRAL ANALYZER AND ITS METHOD			
Patent Number:	EP1167949			
Publication date:	2002-01-02			
Inventor(s):	KIKUGAWA TOMOYUKI (JP); UEHARA KIYOJI (JP); YOSHIDA NAOHIRO (JP)			
Applicant(s):	JAPAN SCIENCE & TECH CORP (JP); ANRITSU CORP (JP)			
Requested Patent:	EP1167949			
Application Number:	EP20000911279 20000322			
Priority Number(s):	WO2000JP01743 20000322; JP19990084898 19990326			
IPC Classification:	G01N21/35; G01N21/39			
EC Classification:	G01N21/31A, G01N21/35B			
Equivalents:	AU3324900,			
Cited Documents:				
Abstract				
An isotopomer absorption spectral analyzing apparatus and its method for precisely measuring the isotope ratio by substantially equalizing the absorption signal levels of different species of isotopes. In an isotopomer absorption spectral analyzing apparatus, a sample cell (21) capable of providing optical paths of different optical lengths is installed, optical beams A, B are caused to enter the sample cell (21) and travel along paths of different optical lengths, thereby determining the abundance ratio between species of isotopes in				
molecules from the ra	atio between intensities of signals corresponding to the species of isotopes.			
Data supplied from the esp@cenet database - I2				

(12)

EUROPEAN PATENT APPLICATION

published in accordance with Art. 158(3) EPC

- (43) Date of publication: 02.01.2002 Bulletin 2002/01
- (21) Application number: 00911279.8
- (22) Date of filing: 22.03.2000

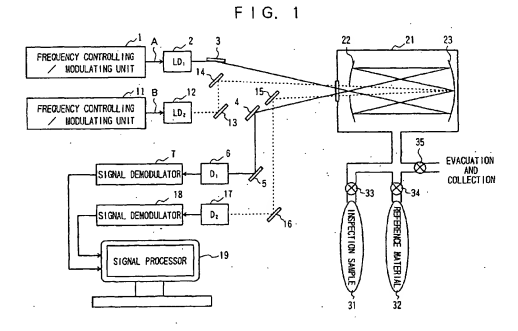
- (51) Int Cl.7: **G01N 21/35**, G01N 21/39
- (86) International application number: PCT/JP00/01743
- (87) International publication number: WO 00/58712 (05.10.2000 Gazette 2000/40)
- (84) Designated Contracting States: AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU MC NL PT SE
- (30) Priority: 26.03.1999 JP 8489899
- (71) Applicants:
 - Japan Science and Technology Corporation Kawaguchi-shi, Saitama (JP)
 - Anritsu Corporation Minato-ku, Tokyo 106-0047 (JP)

- (72) Inventors:
 - UEHARA, Kiyoji Tokyo 135-0033 (JP)
 - YOSHIDA, Naohiro Sagamihara-shi, Kanagawa 228-0803 (JP)
 - KIKUGAWA, Tomoyuki Minato-ku, Tokyo 106-0047 (JP)
- (74) Representative: Howden, Christopher Andrew FORRESTER & BOEHMERT Pettenkoferstrasse 20-22 80336 München (DE)

ISOTOPOMER ABSORPTION SPECTRAL ANALYZER AND ITS METHOD (54)

An isotopomer absorption spectral analyzing apparatus and its method for precisely measuring the isotope ratio by substantially equalizing the absorption signal levels of different species of isotopes. In an isotopomer absorption spectral analyzing apparatus, a sample cell (21) capable of providing optical paths of

different optical lengths is installed, optical beams A, B are caused to enter the sample cell (21) and travel along paths of different optical lengths, thereby determining the abundance ratio between species of isotopes in molecules from the ratio between intensities of signals corresponding to the species of isotopes.



Description

TECHNICAL FIELD

[0001] The present invention relates to an isotopomer absorption spectral analyzing apparatus and method for precisely assaying an isotopomer—a molecule containing an isotope—for inferring the origin thereof, contemplating applications in scientific fields, including environmental analysis; applications in the medical field, including diagnosis; and applications in other fields.

BACKGROUND ART

[0002] Conventional absorption spectral analyzing apparatuses employ a sample cell having a single optical path.

DISCLOSURE OF THE INVENTION

[0003] Therefore, when the isotope abundance ratio (the abundance ratio between two isotopes) deviates greatly from 1:1, a great difference is present between the levels of absorption signals corresponding to the isotope species, depending on the species of the isotopes. For example, in the case of naturally occurring CH₄, the abundance ratio of ¹²CH₄ to ¹³CH₄ is approximately 100:1, and therefore, the absorption signal level of ¹²CH₄ is approximately 100 times that of ¹³CH₄, making precise measurement of the isotope ratio difficult.

[0004] The present invention has been accomplished so as to solve the aforementioned problem. Thus, an object of the present invention is to provide an isotopomer absorption spectral analyzing apparatus and method which enable absorption signals corresponding to different isotopes to assume substantially the same level, to thereby enable precise measurement of the isotope ratio.

[0005] In order to achieve the above objects, the present invention provides the following.

[1] An isotopomer absorption spectral analyzing apparatus, characterized in that a sample cell capable of providing optical paths of different optical lengths is installed; at least two optical beams are caused to enter the sample cell such that the optical beams travel along optical paths of different optical path lengths; and the abundance ratio between species of isotopes in molecules is determined from the ratio between intensities of absorption signals corresponding to the species of isotopes.

[2] An isotopomer absorption spectral analyzing apparatus as described in [1], wherein the at least two optical beams are emitted from a single light source of variable-wavelength type or from a plurality of light sources of fixed-wavelength type or variable-wavelength type.

[3] An isotopomer absorption spectral analyzing ap-

paratus as described in [1], wherein the sample cell is a multiple-reflection absorption cell having paired reflection mirrors.

[4] A method of isotopomer absorption spectral analysis, characterized in that a sample cell capable of providing optical paths of different optical lengths is used in order to substantially equalize levels of absorption signals corresponding to species of isotopes, to thereby enable precise measurement of the abundance ratio between the species of isotopes.

BRIEF DESCRIPTION OF THE DRAWINGS

5 [0006]

FIG. 1 shows the configuration of an isotopomer absorption spectral analysis system according to one embodiment of the present invention.

FIG. 2 shows a modification of the light source used in the isotopomer absorption spectral analysis system according to the embodiment of the present invention.

FIG. 3 shows a modification of the cell used in the isotopomer absorption spectral analysis system according to the embodiment of the present invention.

BEST MODES FOR CARRYING OUT THE INVENTION

[0007] A mode for carrying out the present invention will next be described in detail with reference to the drawings.

[0008] FIG. 1 is shows the configuration of an isotopomer absorption spectral analysis system according to one embodiment of the present invention. The system is based on laser spectroscopy employing a plurality of optical paths.

[0009] In FIG. 1, reference numeral 1 represents a first frequency controlling/modulating unit; reference numeral 2 represents a first laser diode LD₁; reference numerals 3, 4, and 5 represent reflection mirrors; reference numeral 6 represents a first optical detector D₁; reference numeral 7 represents a first signal demodulator; reference numeral 11 represents a second frequency controlling/modulating unit; reference numeral 12 represents a second laser diode LD2; reference numerals 13, 14, 15, and 16 represent reflection mirrors; reference numeral 17 represents a second optical detector D₂; reference numeral 18 represents a second signal demodulator; reference number 19 represents a signal processor; reference numeral 21 represents a long-optical-path cell (sample cell: multiple-reflection absorption cell); reference numerals 22 and 23 represent paired reflection mirrors; reference number 31 represents an inspection sample (in which the abundance ratio of ¹²CH₄ to ¹³CH₄ is unknown); reference number 32 represents a reference material (in which the abundance ratio of ¹²CH₄ to ¹³CH₄ is known); and reference

numerals 33, 34, and 35 represent open-close control valves. In the long-optical-path cell 21, a longer optical path is represented by a continuous line, and a shorter optical path is represented by a dotted line. The optical path length (determined by the number of times of reflection) is set through adjustment of the incident angle of the relevant optical beam and/or the angles of mirrors. [0010] Optical beams A and B from the light sources are caused to enter, at different angles, the long-optical-path cell 21 serving as a multiple-reflection absorption cell and equipped with paired reflection mirrors 22 and 23, so that the optical beams A and B travel along optical paths of different lengths; e.g., 1 m and 100 m. Thus, an optical path difference suited for measuring the abundance ratio of a sample can be provided.

3

[0011] The abundance of an isotopomer (isotopecontaining molecule) varies greatly in accordance with its origin and other factors. Accordingly, through precise measurement of the isotopomer abundance ratio of samples collected from many places in the world, formation, transfer, and disappearance of environmental substances can be analyzed in detail.

[0012] The present invention provides highly effective analysis means which supplements mass analysis conventionally employed for isotopomer analysis. In the method of the present invention, the ratio of the absorption signal of one isotopomer to that of another isotopomer is measured and compared with the same ratio of the reference material 32.

[0013] The ¹³CH₄/¹²CH₄ ratio of methane contained in the atmosphere was measured. The ratio is known to be approximately 1/100. Thus, in order to obtain absolute absorption signals of substantially the same level, there may be employed either one of the following method (1) or (2): method (1) employing combination of a strong incident beam to be absorbed by ¹³CH₄ and a weak incident beam to be absorbed by ¹²CH₄, or method (2) employing two strong beams and causing them to travel along optical paths of different optical path lengths. In the present embodiment, the ratio was measured in accordance with the aforementioned method (2) by use of the method and analyzing apparatus of the present invention equipped with a modified Herriott long-optical-path cell (product of New Focus) and two wavelength-stabilized semiconductor lasers (laser diodes). The provided optical path lengths are 100 m and 1.1 m.

[0014] In the above embodiment, optical beams are generated by use of a plurality of light sources. Alternatively, as shown in FIG. 2, a plurality of optical beams may be generated by use of a single light source (laser diode LD) 42 and a semitransparent mirror for dividing a beam from the light source. In FIG. 2, reference numeral 41 represents a frequency controlling/modulating unit, and reference numerals 44 and 45 represent reflection mirrors. Elements identical to those shown in FIG. 1 are represented by the same reference numerals as shown in FIG. 1, and repeated description thereof is

omitted.

[0015] The aforementioned embodiment uses a cell including reflection mirrors, thereby providing a plurality of optical paths of different optical lengths. However, the structure of the cell is not limited to the aforementioned structure, and a cell as shown in FIG. 3; i.e., a cell having a short optical path portion 51 and a long optical path portion 52, may also be employed.

[0016] The present invention is not limited to the above-described embodiment. Numerous modifications and variations of the present invention are possible in light of the spirit of the present invention, and they are not excluded from the scope of the present invention.

[0017] As described in detail hereinabove, in the present invention, through employment of a sample cell capable of providing a plurality of optical paths of different optical lengths, the levels of absorption signals corresponding to isotope species can be substantially equalized in order to enable precise measurement of the isotope abundance ratio.

INDUSTRIAL APPLICABILITY

[0018] The present invention enables precise assay of an isotopomer—a molecule containing an isotope—for inferring the origin thereof. The invention is expected to find applications in scientific fields, including environmental analysis; and in medical fields including diagnosis.

Claims

35

- An isotopomer absorption spectral analyzing apparatus, characterized in that
 - (a) a sample cell capable of providing optical paths of different optical lengths is installed;
 and
 - (b) at least two optical beams are caused to enter the sample cell such that the optical beams travel along optical paths of different optical path lengths, and the abundance ratio between species of isotopes in molecules is determined from the ratio between intensities of absorption signals corresponding to the species of isotopes.
- An isotopomer absorption spectral analyzing apparatus as described in claim 1, wherein the at least
 two optical beams are emitted from a single light
 source of variable-wavelength type or from a plurality of light sources of fixed-wavelength type or variable-wavelength type.
- An isotopomer absorption spectral analyzing apparatus as described in claim 1, wherein the sample cell is a multiple-reflection absorption cell having

paired reflection mirrors.

4. A method of isotopomer absorption spectral analysis, characterized in that a sample cell capable of providing optical paths of different optical lengths is 5 used in order to substantially equalize levels of absorption signals corresponding to species of isotopes, to thereby enable precise measurement of the abundance ratio between the species of isotopes.

10

15

20

25

30

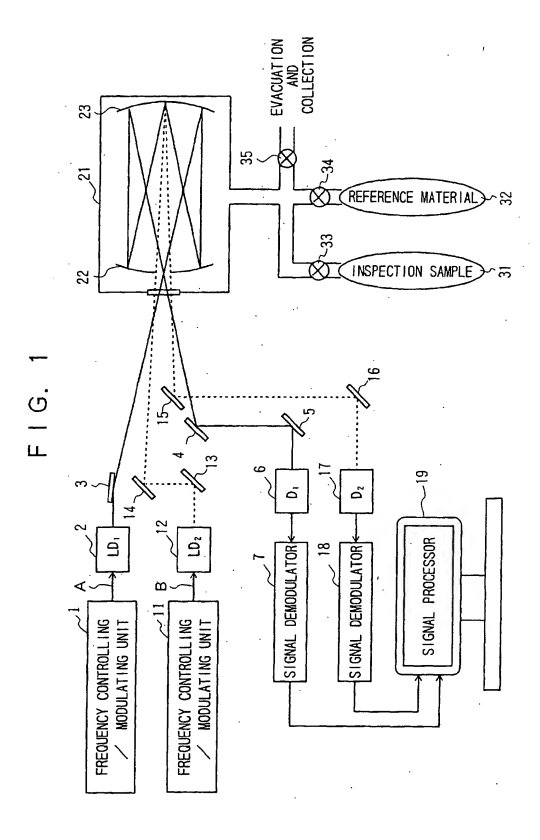
35

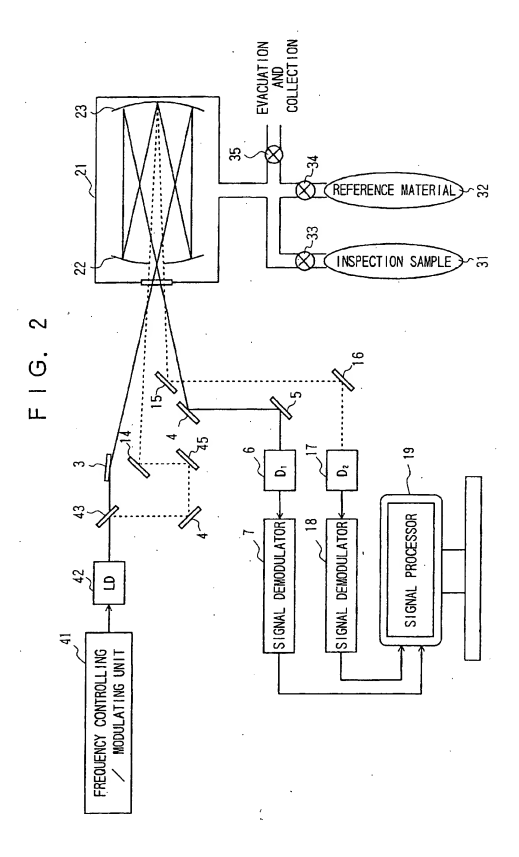
40

45

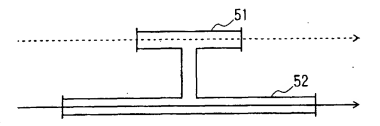
50

55





F I G. 3



INTERNATIONAL SEARCH REPORT

International application No.

PCT/JP00/01743

	SIFICATION OF SUBJECT MATTER . Cl ⁷ G01N21/ 35			
	.Cl ⁷ G01N21/ 39			
According	to International Patent Classification (IPC) or to both i	national classification and IPC		
B. FIELD	S SEARCHED			
	ocumentation searched (classification system follower	d by classification symbols)		
Int	.Cl' G01N21/00-21/61		•	
Documenta	tion searched other than minimum documentation to the	be extent that such documents are included	in the fields searched	
Jits	suyo Shinan Koho 1922-1996	Toroku Jitsuyo Shinan K	loho 1994-2000	
KOKA	i Jitsuyo Shinan Koho 1971-2000	Jitsuyo Shinan Toroku K	oho 1996-2000	
Electronic d	ata base consulted during the international search (nar	me of data base and, where practicable, sea	rch terms used)	
JOIS WPI/				
,,,,,,,	2			
C. DOCU	MENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where a	ppropriate, of the relevant passages	Relevant to claim No.	
Х	US, 5146294, A (Roland Grisar)	,	1-4	
	08 September, 1992 (08.09.92), Full text; Figs. 4 to 5			
	& DE,4012454	İ	-	
х	JP, 10-197444, A (Otsuka Pharm 31 July, 1998 (31.07.98),	aceutical Co., Ltd.),	1,4	
	Full text; Fig. 1	i		
	& WO, 9830888, A & CN, 1243 & AU, 9853439, A & EP, 9531	575, A		
	& AU, 9853439, A & EP, 9531	.48, A		
х	JP, 53-42890, A (Nippon Bunko	Kogyo K.K.),	1,4	
	18 April, 1978 (18.04.78),		·	
	Full text; Fig. 2 (Family: no	one)		
А	JP, 09-297061, A (Japan Radio	Co., Ltd.),	2	
	18 November, 1997 (18.11.97),			
	Full text; Figs. 1, 8 & GB, 2312743, A & DE, 1971	5685 A		
	& US, 5929442			
, [Champanham Champial Marrian) P. 1			
A	Chemosphere Chem Biol Toxicol Rel	at Environ Probl, Vol. 26,	1-4	
Further	documents are listed in the continuation of Box C.	See patent family annex.		
	categories of cited documents: nt defining the general state of the art which is not	"T" later document published after the inter- priority date and not in conflict with the		
consider	ed to be of particular relevance	understand the principle or theory under	rlying the invention	
date	ocument but published on or after the international filing	"X" document of particular relevance; the cl considered novel or cannot be considered		
	nt which may throw doubts on priority claim(s) or which is establish the publication date of another citation or other	step when the document is taken alone "Y" document of particular relevance; the ci	aimed invention cannot be	
special r	eason (as specified) nt referring to an oral disclosure, use, exhibition or other	considered to involve an inventive step	when the document is	
means		combined with one or more other such of combination being obvious to a person	skilled in the art	
"P" documer than the	nt published prior to the international filing date but later priority date claimed	"&" document member of the same patent fa	mily	
	ctual completion of the international search	Date of mailing of the international searc	h report	
13 Jı	une, 2000 (13.06.00)	20 June, 2000 (20.06		
Name and mailing address of the ISA/ Japanese Patent Office		Authorized officer		
oapai	rese sateur Attice			
Facsimile No	exsimile No. Telephone No.			
2000.00				

Form PCT/ISA/210 (second sheet) (July 1992)

INTERNATIONAL SEARCH REPORT

International application No.
PCT/JP00/01743

C (Continua	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant	passages	Relevant to claim
	No.1 (in English), Nuffield Press Ltd, (1993),	pp.13-22	
	•		
		j	
{		1	
		j	
1		1	
		į	
-		- 1	
1		Ī	
į			
- 1			
l			
i		- 1	
- 1			
ļ		1	
}		- 1	
1		ŀ	
1		1	
		1	
İ			
1		1	
		1	
1		}	
		ŀ	
		1	
		ļ	
İ		į	
		İ	
Ì		J	
,		l	
ſ		1	
1		!	
		1	
ĺ		-	
		1	
		ĺ	
		1	
			•
1			
		[
ļ			
ł			
ł			
		1	
		ļ	
- 1			

ç